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201-14854

November 20, 2003

Administrator
US EPA
P.O. Box 1473
Merrifield, VA 22116
Attn: Chemical Right-to-Know Program

OPPT CBIC

Dear Administrator:

On behalf of the member companies of the HPV Committee, the International Association of Color Manufacturers is pleased to submit the test plan and robust summaries for 2-naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt (FD&C Red 40). The IACM HPV Committee has chosen not to belong to the HPV Tracker System for submission of test plans and robust summaries. We are therefore submitting the test plan and accompanying robust summaries directly to EPA to make available to the public. A hard copy of this submission is available upon request. The EPA registration number for the IACM HPV Committee is

Please feel free to contact me with any questions or comments you might have concerning the submission (<u>tadams@therobertsgroup.net</u> or 202-331-2325).

Sincerely,

Timothy Adams, Ph.D.
Technical Contact Person for IACM HPV

Test Plan for 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

OPPT CBIC

Consortium Registration Number

Submitted to the EPA under the HPV Challenge Program by:

The International Association of Color Manufacturers/HPV Committee

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List of Member Companies

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Test Plan for

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

1 IDENTITY OF SUBSTANCES

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Synonyms:

FD&C Red No. 40 Allura Red CI Food Red 17

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2 CATEGORY ANALYSIS

2.1 Introduction

The International Association of Color Manufacturers (IACM) has volunteered to participate in the EPA's Chemical "Right-to-Know" Program. IACM is committed to assembling and reviewing available test data, developing and providing test plans for each of the sponsored chemicals, and, where needed, conducting additional testing on the chemicals used by the color industry in order to assure their human and environmental safety. The category analysis, test plan, and robust summaries presented represent the first phase of IACM's commitment to the Chemical "Right-to-Know" Program.

2.2 BACKGROUND INFORMATION

This category analysis and test plan provides data for FD&C Red No. 40. FD&C Red No. 40 is a red powder that is soluble in water and is used to color gelatins, puddings, custards, alcoholic and nonalcoholic beverages, sauces, topping, candy, sugars, frostings, fruits, juices, dairy products, bakery products, jams, jellies, condiments, meat and poultry. FD&C Red No. 40 is also used to color drugs and cosmetics.

FD&C Red No. 40 is an azo dye. Azo compounds are formed from arenediazonium ions reacting with highly reactive aromatic compounds, in what is called a diazo coupling reaction. Azo compounds are generally deeply colored because the azo linkage brings the two aromatic rings into conjugation [Solomon, 1996]. In addition to possessing extended conjugation, many azo dyes are also ring substituted with sulfonic acid substituents, which significantly increase polarity and water solubility.

2.3 REGULATORY STATUS

FD&C Red No. 40 is a certified color additive approved in the United States to color food, drugs and cosmetics. Certified color additives are synthetic organic compounds that must meet high purity specifications established by the Food and Drug Administration (FDA) (see Table 1 below). Each batch of manufactured certified color in the United States is tested by the FDA for compliance with these specifications [Frick and Meggos, 1988]. Certified color additives are among the most thoroughly studied of all food ingredients because of the rigorous testing for human health endpoints required by the 1960 Color Additive Amendments to the FD&C Act [Hallagan, 1991]. There are currently only seven certified color additives approved for food, drug and cosmetic use in the United States.

Table 1. US FDA Specifications

FD&C Red No. 40 shall conform to the following specifications and shall be free from impurities other than those named to the extent that such other impurities may be avoided by good manufacturing practice (21 CFR 74.340):

- Sum of volatile matter (at 135° C) and chlorides and sulfates (calculated as sodium salts), not more than 14.0 percent.
 - Water-insoluble matter, not more than 0.2 percent.
 - Higher sulfonated subsidiary colors (as sodium salts), not more than 1.0 percent.
 - Lower sulfonated subsidiary colors (as sodium salts), not more than 1.0 percent.
- Disodium salt of 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenoxy)-2-naphthalenesulfonic acid, not more than 1.0 percent.
- Sodium salt of 6-hydroxy-2-naphthalenesulfonic acid (Schaeffer's salt), not more than 0.3 percent.
 - 4-Amino-5-methoxy-o-toluenesulfonic acid, not more than 0.2 percent.
- Disodium salt of 6,6'-oxybis (2-naphthalene-sulfonic acid), not more than 10 parts per million.
 - Lead (as Pb), not more than 10 parts per million.
 - Arsenic (as As), not more than 3 parts per million.
 - Total color, not less than 85.0 percent.

FD&C Red No. 40 was first listed for food use in the United States in 1971. In 1994, 1,477,651 kg of FD&C Red No. 40 dye and 257,752 kg of FD&C Red No. 40 lake were certified for use in the United States.

The World Health Organization/Food and Agriculture Organization Joint Expert Committee for the Evaluation of Food Additives (WHO/FAO JECFA) has also evaluated the safety of FD&C Red No. 40 used as a coloring agent in food. An average daily intake (ADI) of 0-7 mg/kg bw/d was assigned by JECFA in 1981 based on the extensive human toxicological information available that indicated FD&C Red No. 40 did not possess carcinogenic potential (see Table 2 below).

Table 2. Regulatory Approvals/Consumption Limits ¹			
USA EEC	GMP (21 CFR 74.340) GMP (EC Journal No. L237; 1994)		
JECFA	ADI of 0-7 mg/kg body weight (25th Report, 1981)		

Based on the long history of use of FD&C Red No. 40 in food, the many hazard assessments performed by the United States FDA and WHO/FAO JECFA, and the current regulatory status of FD&C Red No. 40, there is no compelling evidence that this substance should be further tested for human health endpoints in the EPA Chemical "Right to Know" Program.

2.4 STRUCTURAL CLASSIFICATION

FD&C Red No. 40 is principally the disodium salt of 6-hydroxy-5-[(2-methoxy-5-methyl-4sulfophenyl) azo]-2-naphthalenesulfonic acid (US FDA-21 CFR 74.340). FD&C Red No. 40 is a monoazo dye. The diazo nucleus (-N=N-) contains a benzene ring

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¹ IACM, 2003

substituted with *o*-methoxy, *m*-methyl, and *p*-sulfonic acid groups and a naphthalene ring substituted with *o*-hydroxy and *p*'-sulfonic acid groups.

2.5 Industrial Production

FD&C Red No. 40 is manufactured by coupling diazotized 5-amino-4-methoxy-2-toluenesulfonic acid with 6-hydroxy-2-naphthalene sulfonic acid.

2.6 PHARMACOKINETICS AND METABOLISM

Orally administered FD&C Red No. 40 is poorly absorbed by dogs and rats. Dogs (2) were pretreated with unlabeled 100 mg/kg bw/d FD&C Red No. 40 in capsules for five days and with 100 mg/kg bw/d ³⁵S radiolabelled FD&C Red No. 40 on the sixth day. The animals were studied for up to 72 hours following the last dose. After 72 hours, no significant radioactivity could be detected in any of the organs and tissues assayed. The major excretory route was through the feces. Approximately 85% of the administered dose (75 and 95% in each) was excreted in the feces within 24 hours. Minor amounts (1.1 and 0.3%) were excreted during the following 24-48 hour collection period. The urinary excretion of FD&C Red No. 40 was minimal with an average recovery of 3% in dogs. The total recovery of radioactivity from all tissues assayed was insignificant, with the exception of the intestine and its contents [Hazelton Laboratories, 1975a].

Five male rats were pretreated with unlabeled 200 mg/kg bw/d FD&C Red No. 40 *via* gavage for five days and with 200 mg/kg bw/d ³⁵S radiolabelled FD&C Red No. 40 on the sixth day. Two controls received the vehicle only. The animals were studied for up to 72 hours following the last dose. Similar to dogs, the majority of the radioactivity was excreted in the feces (76-92% in 72 hours) with minimal amounts excreted in the urine (approximately 8%²) as the parent substance or its metabolites [Hazelton Laboratories, 1975a].

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² 19.8% of the radioactivity administered was excreted in the urine in one test animal, but given the value was nearly double that of any other animal, and the fact that lung damage was observed at necropsy, the authors reported the animals was improperly dosed and that the value was excessive.

In a separate follow-up study in both rats and dogs using the protocol described above, the principal urinary metabolite in both species was determined to be 4-amino-5-methoxy-o-toluene sulfonic acid (cresidine sulphonic acid), indicating diazo reduction and cleavage to yield two amine fragments. Cresidine sulphonic acid was also the principal urinary metabolite (greater than 90%). However, total urinary metabolites only accounted for approximately 7% of the total administered dose. Analysis of the rat-fecal extracts revealed an unknown metabolite (35.4%); unchanged FD&C Red No. 40 at (14.9%); and cresidine sulphonic acid (25.6%). The formation of cresidine sulfonic acid in the feces may arise *via* microfloral reduction of the parent dye. A similar metabolic pattern was reported following the analysis of the dog-fecal extract. The authors commented that the polar unknown metabolite is apparently a glucuronide or sulfate conjugate, possibly formed from one of the two diazo reduction products, 1-amino-2-naphthol-6-sulfonic acid. The polar unknown is not absorbed, given that only 0.6% of the dose was found at the origin following thin layer chromatography analysis of the rat urine [Hazleton Laboratories, 1975b].

In summary, FD&C Red No. 40 is predominantly not absorbed by animals. It is excreted mainly in the feces either unchanged or as polar metabolites (*e.g.*, cresidine sulfonic acid) formed by diazo reduction FD&C Red No. 40. Fecal excretion occurs predominantly within the first 24 hours. Small amounts of cresidine sulfonic acid are also detected in the urine.

3 TEST PLAN

3.1 CHEMICAL AND PHYSICAL PROPERTIES

3.1.1 Melting Point

FD&C Red No. 40 is a solid and did not melt when heated to 300 °C [Hazelton Laboratories, 1970]. Accordingly, the melting point of FD&C Red No. 40 was calculated to be 350 °C using modeling software [MPBPVPWIN EPI Suite, 2000].

3.1.2 Boiling Point

The boiling point of FD&C Red No. 40 was calculated to be 872 °C [MPBPVPWIN EPI Suite, 2000]. Technically, data for this endpoint are not required given that this material is a solid and would likely decompose upon heating to elevated temperatures.

3.1.3 Vapor Pressure

The calculated vapor pressure for FD&C Red No. 40 has been reported to be 1.25 X 10⁻²³ mm Hg at 25°C [MPBPVPWIN EPI Suite, 2000]. Given the high molecular mass of FD&C Red No. 40 (496.43) and the estimated Henry's law constant for azo dyes of 10⁻¹⁵ atm-m³/mol it is highly unlikely that FD&C Red No. 40 would exhibit any significant (less than 0.001 mm Hg) vapor pressure. This is predicted by the MPBPVPWIN model. Based on these data, the vapor pressure is less than 1 X 10⁻²⁰ mm Hg.

3.1.4 Octanol/Water Partition Coefficients

Log K_{OW} value for FD&C Red No. 40 is -0.55 [KOWWIN EPI Suite, 2000]. The experimental log Kow value would be difficult to obtain by OECD methods given the large difference between water solubility and anticipated solubility in octanol. Based on the observations that FD&C Red No. 40 is very water soluble (220,000 mg/L) and

essentially insoluble in a relatively polar solvent like ethanol (1 mg/L) [Marmion, 1991], it is anticipated that the log Kow value for this substances would exceed 6.0.

3.1.5 Water Solubility

FD&C Red No. 40 has a reported water solubility of 180,000 mg/L at 20 °C, 220,000 mg/L at 25 °C, and 260,000 mg/L at 60 °C [Marmion, 1991]. The solubility of FD&C Red No. 40 in 100% glycerol is 3,000 mg/L at 25 °C while the solubility in ethanol is reported to be 1 mg/L at 25 °C [Marmion, 1991, robust summary not included]. The solubility of FD&C Red No. 40 in octanol is expected to be less than 1 mg/L.

3.1.6 New Testing Required

None.

3.2 ENVIRONMENTAL FATE AND PATHWAYS

3.2.1 Photodegradation

Direct and indirect photolysis experiments were conducted on FD&C Red No. 40 using two 15-watt low pressure lamps as the ultraviolet light source. Following 50 minutes of exposure to the lamps, FD&C Red No. 40 concentration decreased by 7% in the direct experiment. In the indirect experiment which used acetone as the sensitizer, the concentration of FD&C Red No. 40 decreased by 99% after 20 minutes [Pasin and Rickbaugh, 1991]. The calculated half-life for hydroxyl radical reactions is 18.2 hours [AOPWIN EPI Suite, 2000].

3.2.2 Stability In Water

FD &C Red No. 40 does not contain functional groups (*e.g.*, esters, amides, acetals, epoxides, lactones, *etc.*) that hydrolyze in water. The only potential reactivity in water would involve desulfonation of the aromatic sulfonic acid or its corresponding sulfonic acid salt. In aqueous acid (sulfuric acid), aromatic sulfonic acids desulfonate at temperatures of 100 to 175 °C. These conditions would not typically be encountered in the environment. Therefore, FD &C Red No. 40 and its corresponding salts are anticipated to be stable in water.

3.2.3 Biodegradation

The biodegradability of azo dyes substituted with a phenolic OH and two sulfonic acid groups consistently show that these substances are not absorbed onto activated sludge and, therefore, are not biodegradable [Shaul *et al.*, 1990]. Incubation of 1.0 or 5.0 mg/L of a structurally related azo dye, (1-naphthalenesulfonic acid, 4-hydroxy-3-[(4-sulfo-1-

naphthalenyl)azo]-, disodium salt)³ with activated sludge from a sewage treatment plant revealed that the concentration of dye remained essentially constant in the influent flow, primary effluent, and activated sludge effluent. Essentially no azo dye was absorbed by activated sludge. Two other azo dyes ring-substituted with sulfonic acid groups (Acid Orange No. 10 and Acid Red No. 1) exhibited a similar behavior in these experiments.

FD&C Red No. 40 was not predicted to be readily degradable by BIOWIN model calculations [AOPWIN EPI Suite, 2000].

3.2.4 Fugacity

Transport and distribution in the environment were modeled using Level III Fugacity-based Environmental Equilibrium Partitioning Model Version 2.70 [Trent University, 2002]. The principal input parameters into the model are molecular weight, melting point, vapor pressure, water solubility, and log K_{OW}.

As expected, the model predicts that FD&C Red No. 40 is distributed completely to the water compartment (greater than 100%). Consistent with the extremely high water solubility and low log K_{OW} data, FD &C Red No. 40 showed no significant distribution to the soil compartment (2 x 10⁻¹⁴%). Based on this physiochemical model, the ratio for distribution of FD&C Red No. 40 between water (greater than 100%) and fish (4.9 x 10⁻⁶%) is greater than seven orders of magnitude suggesting essentially no bioaccumulation in fish. These data are consistent with ecotoxicity data for aromatic sulfonic acid derivatives that demonstrate essentially no absorption and toxicity to fish even at concentrations exceeding 1000 mg/L.

3.2.5 New Testing Required

None.

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3.3 ECOTOXICITY

3.3.1 Acute Toxicity to Fish

Based on input parameters for molecular weight (496.43), water solubility (220,000 mg/L at 25 °C), and melting point (350 °C), the calculated 96-hour LC50 for FD&C Red No. 40 is 2,714 mg/L [ECOSAR EPI Suite, 2000] indicating a very low order of acute toxicity. The extensive water solubility and limited lipophilicity of FD&C Red No. 40 is to a large extent, a function of the presence of aromatic sulfonic acid and phenolic ring substituents. The extensive studies on the ecotoxicity of aromatic sulfonic acids indicate a very low order of toxicity to fish [Greim *et al.*, 1994]. Experimental LC50 values are available for stilbene sulfonic acids in which the N atom in the diazo dye is replaced by C. As indicated in Table 3 below, acute fish toxicity studies on salts of stilbene sulfonic acid derivatives result in an 96-hour LC50 value greater than 10,000 mg/L. Also, 48-hour and 72-hour LC50 concentrations of 200 and greater than 1000 mg/L, respectively have been reported [Greim *et al.*, 1994]. These values are consistent with calculated values.

Table 3

Name Acute Toxicity to fish

2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid

2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid, disodium salt

72-hour LC50: greater than 1000 mg/L

48-hour LC50: 200 mg/L

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2,2'-(1,2-ethene-diyl)bis(5-amino)benzenesulfonic acid, dipotassium salt

96-hour LC50: greater than 10,000 mg/L

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Given the high-calculated LC50 values from the ECOSAR model, the experimentally measured toxicity of aromatic sulfonic acid derivatives, and the difficulties inherent in acute aquatic testing with dyes, no additional testing is requested.

3.3.2 Acute Toxicity to Aquatic Invertebrates

The calculated 48-hour LC50 value for FD&C Red No. 40 in daphnids is 295 mg/L based on input parameters for molecular weight (496.43), water solubility (220,000 mg/L at 25 °C), and melting point (350 °C), [ECOSAR EPI Suite, 2000] indicating a low order of acute toxicity. The extensive water solubility and limited lipophilicity of FD&C Red No. 40 is to a large extent, a function of the presence of aromatic sulfonic acid phenolic ring substituents. The extensive studies on the ecotoxicity of aromatic sulfonic acids indicate a very low order of toxicity to aquatic invertebrates [Greim *et al.*, 1994]. An experimental 24-hour EC50 value with *Daphnia* for a stilbene sulfonic acid derivative, 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid, was greater than 100 mg/L [Greim *et al.*, 1994]. This value is consistent with calculated values.

3.3.3 Acute Toxicity to Aquatic Plants

Based on input parameters for molecular weight (496.43), water solubility (220,000 mg/L at 25 °C), and melting point (350 °C), the calculated EC50 for FD&C Red No. 40 with

green algae is 44,524 mg/L [ECOSAR EPI Suite, 2000] indicating a very low order of acute toxicity. In a 96-hour algal chronic toxicity test, a sulfonic acid substituted azo dye, stimulated population growth (26.4%) compared to control (algal assay medium) [Greene and Baughman, 1996]. In fact, of the 46 dyes tested, only one, an anthraquinone dye, produced measurable toxicity. Given the high-predicted value for acute toxicity to aquatic plants and the stimulation of plant growth resulting from the addition of a structurally related azo dye in an experimental acute toxicity test, it is not recommended that additional tests be performed.

3.3.4 New Testing Required

None.

3.4 HUMAN HEALTH TOXICITY

3.4.1 Acute Toxicity

Pre-GLP acute toxicity studies were conducted on FD&C Red No. 40 in rats and dogs. Six groups of five male and five female Sprague-Dawley rats were each administered the test substance in a 10% weight/volume solution. The dosage levels tested were 215, 464, 1,000, 2,150, 4640, and 10,000 mg/kg bw. Observations were made immediately following dosing, at 1, 4, 24, 48-hours and once daily thereafter up to 14 days. Following the observation period, the animals were weighed, sacrificed by cerebral concussion and necropsied. Clinical observations were normal with the exception of red-colored feces in both sexes at all dose levels and red-colored urine at the three highest dose levels in the female animals. There were no deaths at any dose level tested. The acute LD50 was determined to be greater than 10,000 mg/kg bw/day for adult male and female Sprague-Dawley albino rats administered FD&C Red No. 40 *via* gavage [Hazelton Laboratories, Inc., 1965a].

Two male Mongrel dogs were administered the test substance in an aqueous solution at a dose level of 5,000 mg/kg bw. Observations were made immediately following dosing and daily thereafter for 7 days. Following the observation period, the animals were weighed, sacrificed and necropsied. Red diarrhea was observed 30 minutes following dosing in one animal, which was followed by emesis. Red urine was reported for the other animal. Red stools were reported for both dogs one day following dosing. From the third day until the seventh day, both animals appeared normal with respect to appearance, behavior, appetite and elimination. Gross necropsy revealed fibrotic changes and decreased weight in a kidney of one test animal. This finding was not considered treatment related, but was rather considered to be a chronic lesion. The spleen also appeared enlarged in this test animal. In the other test animal, hookworms were observed in the gastrointestinal tract. There were no deaths at the dose level tested (5,000 mg/kg bw). The acute LD50 was determined to be greater than 5,000 mg/kg bw/day for male

Mongrel dogs administered FD&C Red No. 40 *via* gavage [Hazelton Laboratories, Inc. 1965b].

3.4.2 In vitro and In vivo Genotoxicity

3.4.2.1 In vitro

FD&C Red No. 40 tested negative in reverse mutation assay using TA1535, TA1537, TA98, TA100; and *Saccharomyces cerevisiae* strain D4 with and without metabolic activation at concentrations up to 5,000 micrograms/plate [Brusick, 1976; Muzzall and Cook, 1979].

3.4.2.2 In vivo

In vivo genotoxicity data are available for the structurally related azo dye, FD&C Yellow No. 6 (6-hydroxy-5-[(4-sulfophenyl)azo]-2-napthalenesulfonic acid, disodium salt)⁴, which is the same structure as FD&C Red No. 40, except without the 2-methoxy and 5-methyl functional groups on the single ring. FD&C Yellow No. 6 tested negative in the rat micronucleus test at a single dose level of up to 1,000 mg/kg bw/day [Westmoreland and Gatehouse, 1991].

3.4.3 Repeat Dose Toxicity

In a Lifetime Toxicity/Carcinogenicity Study, FD&C Red No. 40 was provided in the diet as an admixture to Sprague-Dawley rats. In the *in utero* phase, 240 male and female rats were randomly assigned (30/group) to the control, low dose (0.37%), mid-dose (1.39%) or high dose (5.19%) groups, providing daily intake levels of 180, 701 or 2,829

NaO₃S
$$\sim$$
 N=N \sim SO₃Na

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mg/kg bw/day for males and 228, 901 or 3,604 mg/kg bw/day for females. These parental (P₁) rats received the test material one week prior to mating, during the three-week mating period and during the gestation and lactation periods. The offspring of these animals were randomly selected and put into groups of fifty male and female weanling rats each. These groups were administered the test substance in the diet of the male animals for 118 weeks and the diet of female animals for 121 weeks at levels of 0, 0.37, 1.39 to 5.19 % corresponding to the dietary levels used in the *in utero* phase. Parameters included survival, clinical signs, body weight and food consumption, gross and microscopic pathology. Gross necropsies were performed on all animals dying during the study, all animals found in a moribund condition, and all animals killed at study termination. Complete histological examinations were performed on all animals in both the control and high-dose groups. The tissues examined histologically included: brain, pituitary, thoracic spinal cord, eyes, esophagus, thyroid, thymus, heart, lungs, liver, spleen, pancreas, stomach, small and large intestine, mesenteric lymph node, kidneys, adrenal, urinary bladder, uterus, prostate, ovaries, testes with epididymides, seminal vesicles, skin, rib junction, bone marrow, nerve with muscle, and any tissue masses or lesions. Histological examination was also performed on animals from any group with observable masses or lesions. If a potential effect was seen recurrently in a tissue, than that tissue was examined in all animals.

Food consumption was elevated among high dose males and females, but was not statistically significant. Red-tinted fur was reported among all treated animals, and red-tinted feces were reported for mid- and high-dose male and females. Group mean body weights of treated males and females were decreased compared to control animals at study termination, with the exception of mid-dose treated male rats that experienced an increase in mean body weight. However, the decrease in mean body weight was only statistically significant in female rats at the high dose level (3,604 mg/kg bw/day). Clinical chemistry and urinalysis parameters revealed no treatment related effects.

Histopathological examination revealed lesions in both control and treated animals at similar prevalence, and thus not attributed to test substance administration. No biologically significant adverse effects were reported following administration of FD&C

Red No. 40, with the exception of decrease mean body weights for high-dose female rats at study termination. The authors attributed this effect to the large amount of non-nutritive material in the diet at the intake level [Borzelleca *et al.*, 1991a].

A similar lifetime/carcinogenicity study was also performed in Charles River HaM/ICR (CD-1) mice and in CD-1 outbred mice. In the *in utero* phase, 50 male and female CD-1 mice each (study A) or 70 male and female CD-1 outbred mice each (study B) were randomly assigned to the control, low dose (0.37%), mid-dose (1.39%) or high dose (5.19%) groups, providing daily intake levels of 507, 1,877 or 7,422 mg/kg bw/day for males and 577, 2,043 or 8,304 mg/kg bw/day for females (study A) and 492, 1,821, or 7,318 mg/kg bw/day (males) and 526, 2,057 or 8,356 mg/kg bw/day (females) (study B). These F₀ groups received the test material one week prior to mating, during the three week mating period and during gestation and lactation periods.

Groups of fifty male and female weanling CD-1 albino mice were randomly selected from the litters at 21 days of age and administered the FD&C Red No. 40 in the diet of study A animals for 104 weeks and the diet of study B animals for 109 weeks at levels of 0, 0.37, 1.39 or 5.19 %. These animals were the F_1 offspring of parental rats (F_0), which were treated at the corresponding levels. Study A had one control group while study B had two control groups. Parameters included survival, clinical signs, body weight and food consumption, gross and microscopic pathology. Gross necropsies were performed on all animals dying during the study, all animals found in a moribund condition, and all animals killed at study termination. Complete histology was conducted on all mice from all groups in study A and on 10/sex/group for the two control groups and the highestdose group from study B. The tissues examined histologically included: brain, pituitary, thoracic spinal cord, eyes, esophagus, thyroid, thymus, heart, lungs, liver, spleen, pancreas, stomach, small and large intestine, mammary glands (study B only), mesenteric lymph node, kidneys, adrenal, urinary bladder, uterus, prostate, ovaries, testes with epididymides, seminal vesicles, skin, rib junction, bone marrow, nerve with muscle, and any tissue masses or lesions. Histological examination was also performed on animals from any group with observable masses or lesions. If a potential effect was seen recurrently in a tissue, than that tissue was examined in all animals.

No treatment-related effects on survival were found. The authors reported decreased food consumption among the mid- and high-dose females for week 62-106 in study B. However, no consistent statistically significant effects on food consumption were reported in either study. Localized alopecia, labored respiration, colored hair coat, lacrimation and thinness were reported in similar incidences in both control and treated mice at all dose levels. Distended abdomens were noted in both mid- and high-dose females, while palpable masses were reported in control and treated groups at a similar incidence. Hematological and clinical chemistry parameters revealed few differences among treated and control groups. No significant gross pathological changes were reported among treated groups compared to control groups. An increase in absolute and relative thyroid weights in study B in the high-dose males and females was reported, but the significance was questioned because there was no accompanying histopathology, nor was it dose-dependent and it appeared to be species-specific.

The authors reported an earlier appearance of lymphatic lymphomas among treated groups in study A compared to control groups. No increases in incidence or appearance of lymphocytic lymphomas were reported in study B. The authors noted that study B was conducted using a different strain of mouse to further investigate if FD&C Red No. 40 had an effect on the appearance of lymphocytic lymphomas, and it revealed no relationship between the incidence of lymphocytic lymphomas and FD&C Red No. 40 [Borzelleca *et al.*, 1991b].

3.4.4 Developmental Toxicity

Four groups of female Osborne-Mendel (FDA strain) rats (40-41 per group) were administered FD&C Red No. 40 in the drinking water at intake levels of 0, 0.2, 0.4 or 0.7% for the first 20 days of gestation. These intake levels correspond to daily doses of 0, 273.58, 545.68 or 939.29 mg/kg bw/day [Collins *et al.*, 1989a]. On day 20, the animals were examined for gross abnormalities followed by euthanasia. Caesarean sections were performed. The uterus was examined for presence and position of resorption sites and fetuses, number of *corpora lutea* and implantation sites. All live fetuses were promptly weighed, sexed, and examined. Crown-rump lengths were measured. Fetuses were

divided and assigned to skeletal or soft tissue examination. No clinical findings were reported and no deaths occurred during treatment. Mean fluid consumption was significantly increased in animals at the 0.2 and 0.4% intake levels, but only on days 14-20. Because fluid consumption was not increased at the 0.7% level, the findings were not considered significant. No other effects were reported.

A significant increase in the incidence of litters containing fetuses with missing sternebrae occurred in the 0.4% group, but not in the group receiving 0.7%. No dose related increases were reported for any sternebral variations. The number of fetuses with at least one type of sternebral variations was greater in all treated groups, but only significantly greater in the 0.4 and 0.7% groups. The percentage of total fetuses with at least one sternebral variation was greater in all of the treated groups compared to the control group, but the differences were not significant. The number of fetuses with more than one skeletal variation were similar among treated and control groups. The incidence of reduced ossification of the hyoid bone was significantly increased at the 0.7% intake level. Significant dose related increases were reported at the highest intake level for the average number of fetuses per litter with at least two skeletal variations and the number of litters containing them.

The authors questioned the biological significance of the reduced ossification of the hyoid bone given the lack of effect seen in a gavage study using higher dose levels. The increased incidence was slightly above that found in the historical controls, and the control group was noted as having a lower incidence compared to the historical controls [Collins *et al.*, 1989a].

Four groups of female Osborne-Mendel (FDA strain) rats (42-43 per group) were administered FD & C Red No. 40 *via* gavage at dose levels of 0, 30, 75, 150, 300, 600 or 1000 mg/kg bw/day for the first 19 days of gestation. On day 19, the animals were examined for gross abnormalities followed by euthanization. Caesarean sections were performed. The uterus was examined for presence and position of resorption sites and fetuses, number of corpora lutea and implantation sites. All live fetuses were promptly

weighed, sexed, and examined. Crown-rump lengths were measured. Fetuses were divided and assigned to skeletal or soft tissue examination.

No clinical findings were reported and no deaths occurred during treatment. No other dose related findings were reported. The only significant skeletal anomaly found was an increase in 14th rib buds at the 300 mg/kg bw/day dose level but was not seen at the higher dose levels. No other soft-tissue or sternebral variations were reported. The NOAEL's for maternal and fetal toxicity were 1000 mg/kg bw/day [Collins *et al.*, 1989b].

3.4.5 Reproductive Toxicity

Groups of male (10) and female (20) Charles River rats were administered FD&C Red No. 40 in the diet at 0, 3700, 13,900, or 51,900 ppm for 27 weeks prior to initiation of the first breeding phase. This P₁ parental generation was individually housed. Clinical observations included food consumption, appearance, individual body weights and behavior and were made weekly.

During the breeding phase of the P₁ generation, two females and one male were placed in a breeding cage. At weekly intervals during the mating period, the males were rotated among the females in each group. Following mating, the females were placed in individual cages to produce the first (FIA) litters. Twenty-four hours following the birth of the pups the first litters (FIA) were arbitrarily reduced to 8 maximum per mother. The number of conceptions, number of litters, live births, stillbirths, size of natural and nursing litters, deaths during the period of lactation, and number of pups weaned were recorded. The body weights of each pup were recorded at 24 hours and at weaning. Gross signs of toxicity were monitored. After 21 days of nursing, random pups were sacrificed and gross necropsies performed. Twenty-four females and twelve males remaining from each test group and control group were selected at random and designated the P₂ generation. Following the weaning of the F1A animals, the P₁ generation was remated to produce their second litters referred to as F1B, according to the procedures described above.

The P₂ generation was housed 4-5 per cage and was maintained on the same dietary levels as their parents. The procedures outlined above for the P₁ generation were maintained for the P₂ generation. The litters of the P₂ animals were referred to as the F2A litters. Body weights of the F2A pups were monitored 24 hours following the birth and at weaning. Gross signs of toxicity were recorded. Following a 21 day nursing period, all pups were weaned and sacrificed. One week following the weaning period of the F2A litter, the P₂ generation was remated to produce their second litters (F2B). Two females were placed in a cage with a male from the corresponding dose group. Males were rotated weekly, and females were examined daily for presence of spermatozoa for a maximum of 21 consecutive days. The first day that sperm were observed was designated as day 0 of gestation. The females were then placed in individual cages. Half of the females (12) were sacrificed on day 19 or 20 of gestation and Caesarean sections were performed. Observations included number and placement of implantation sites, resorption sites, and live and dead fetuses, individual fetal weight and length (crown to rump), and external fetal anatomical structure. Gross necropsies were performed on each female including examination of uterus and visceral structures. The remaining 12 females were allowed to litter normally. The fetuses of both females delivering normally and via Caesarean section were necropsied.

Fertility indices for the control and test animals of both F1A and F1B were considered low. The authors attributed this to the advanced age of the animals upon mating. The fertility index of the 3,700 ppm test group in the F2A breeding cycle as well as the 3700 and 51,900 ppm test groups in the F2B breeding cycle were reported to be low in comparison to control animals and historical control data. Growth suppression, characterized as slight, was also reported for the low-level F1B pups, and the high-level F1A and F1B pups and the F2A and F2B breeding cycles when compared with controls. All other measured parameters were comparable to controls in each generation and among the two filial generations. The authors concluded that FD&C Red No. 40 caused meaningful growth suppression in the pups whose parents received the high level diets. The authors reported a no observable adverse effect level (NOAEL) for reproductive

toxicity following administration of FD&C Red No. 40 as 13,900 ppm [Hazelton Laboratories, 1969].

3.4.6 New Testing Required

None.

3.5 TEST PLAN TABLE

	Physical-Chemical Properties						
Chemical	Melting Poi	nı ı	iling oint	Vapor Pressure	Partition Coefficient	Water Solubility	
2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2- methoxy-5-methyl-4- sulfophenyl)azo]-, disodium salt CAS No. 25956-17-6	A	Calc		Calc	Calc	A	
	Environmental Fate and Pathways						
Chemical	Photodegradation		Stability Water		legradation	Fugacity	
2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2- methoxy-5-methyl-4- sulfophenyl)azo]-, disodium salt CAS No. 25956-17-6	A, Calc		NA		R	Calc	
	Ecotoxicity						
Chemical	Acute Toxicity to Fish		Acute Toxicity to Aquatic Invertebrates		Acute Toxicity to Aquatic Plants		
2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2- methoxy-5-methyl-4- sulfophenyl)azo]-, disodium salt CAS No. 25956-17-6	R, Calc		R, Calc		R, Calc		
	Human Health Data						
Chemical	Acute Toxicity	Genetic Toxicity In Vitro	Genetic Toxicity In Vivo	Repeat Dose Toxicity	Repro- ductive Toxicity	Develop- mental Toxicity	
2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2- methoxy-5-methyl-4- sulfophenyl)azo]-, disodium salt CAS No. 25956-17-6	A	A	R	A	A	A	

Legend			
Symbol	Description		
R	Endpoint requirement fulfilled using category approach, SAR		
Test	Endpoint requirements to be fulfilled with testing		
Calc	Endpoint requirement fulfilled based on calculated data		
A	Endpoint requirement fulfilled with adequate existing data		
NR	Not required per the OECD SIDS guidance		
NA	Not applicable due to physical/chemical properties		

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Robust Summaries for 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Consortium Registration Number

OPPT CBIC

Submitted to the EPA under the HPV Challenge Program by:
The International Association of Color Manufacturers/HPV Committee
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Robust Summaries for

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

The evaluation of the quality of the following data uses a systematic approach described by Klimisch [Klimisch *et al.*, 1996]. Based on criteria relating to international testing standards for categorizing data reliability, four reliability categories have been established. The following categories are:

Reliability code 1. Reliable without restrictions
Reliable with restrictions

Reliability code 3. Not reliableReliability code 4. Not assignable

1 CHEMICAL AND PHYSICAL PROPERTIES

1.1 MELTING POINT

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt
CAS No.	25956-17-6
Method/guideline	Measured
GLP	No
Year	1970
Decomposition	300 °C
Remarks for Results	Decomposes without melting.
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
References	Hazelton Laboratories (1970) Petition to FDA.

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt	
CAS No.	25956-17-6	
Method/guideline	Calculated	
Melting Point	350 °C	
Data Qualities Reliabilities	Reliability code 4. Not assignable.	
Remarks for Data Reliability	Code 4. Calculated.	
References	MPBPVPWIN EPI Suite (2000) US Environmental Protection Agency.	

1.2 BOILING POINT

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt	
CAS No.	25956-17-6	
Method/guideline	Calculated	
Boiling Point	872 °C	
Data Qualities Reliabilities	Reliability code 4. Not assignable.	
Remarks for Data Reliability	Code 4. Calculated.	
References	MPBPVPWIN EPI Suite (2000) US Environmental Protection Agency.	

1.3 VAPOR PRESSURE

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-			
	4-sulfophenyl)azo]-, disodium salt			

CAS No. 25956-17-6

Method/guideline Calculated/Mean of Antoine & Grain

Vapor Pressure 1.25 X 10⁻²³ mm Hg

Temperature 25 °C

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References MPBPVPWIN EPI Suite (2000) US Environmental Protection

Agency.

1.4 N-OCTANOL/WATER PARTITION COEFFICIENTS

Agency.

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-	
CAS No.	4-sulfophenyl)azo]-, disodium salt 25956-17-6	
Method/guideline	Calculated	
Log Pow	-0.55	
Data Qualities Reliabilities	Reliability code 4. Not assignable.	
Remarks for Data Reliability	Code 4. Calculated.	
References	KOWWIN EPI Suite (2000) US Environmental Protection	

1.5 WATER SOLUBILITY

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt			
CAS No.	25956-17-6			
Remarks for Substance	Purity not listed			
Method/Guideline	Experimental			
GLP	Ambiguous			
Year	1991			
Value (mg/L) at Temperature	180,000 mg/L at 20 °C; 220,000 mg/L at 25 °C; 260,000 mg/L at 60 °C			
Description of Solubility	Not given			
Data Qualities Reliabilities	Reliability code 4. Not assignable.			
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).			
References	Marmion D.M. (1991) Handbook of U.S. Colorants: Foods, Drugs, and Cosmetics and Medical Devices. 3rd Ed. New York, John Wiley & Sons, Inc.			

2 ENVIRONMENTAL FATE AND PATHWAYS

2.1 PHOTODEGRADATION

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance FD&C Red No. 40

Method/guideline Not given

Test Type Experimental

GLP Ambiguous

Year 1991

Light Source 15-watt General Electric germicidal lamps

Light Spectrum (nm) Ultraviolet

Remarks for Test Conditions The concentration of the dye solution was measured before

and after the photolysis using the Hewlett-Packard 8452A diode-array UV/Visible Spectrophotometer. FD & C Red No. 40 was prepared in an initial concentration of 5 mg/l. In the first part of the study, photolysis experiments were conducted using two 15-W (30 Watts total) General Electric germicidal lamps as the ultraviolet light source. The distance between the light source and the reaction vessels was approximately 2.5 cm. Both direct photolysis and indirect photolysis experiments were conducted. The indirect photolysis experiment used acetone as

the sensitizer for indirect photodegradation.

Concentration of Substance 5 mg/L

Direct photolysis 7% degradation after 50 minutes

Indirect photolysis 99% degradation after 20 minutes

Sensitizer Acetone

Concentration of sensitizer 5 mg/L

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Pasin B. and Rickabaugh J. (1991) Destruction of Azo Dyes by

Sensitized Photolysis. Hazard. Ind. Wastes, 359-367.

Substance Name

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Method/guideline Calculation

Test Type AOPWIN

Halflife t1/2 18.2 hours

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References AOPWIN EPI Suite (2000) US Environmental Protection

Agency.

2.2 BIODEGRADATION

Substance Name	2-1	Naph	nthale	enes	ulfo	nic a	acid,	6-hydroxy-5-[(2-methoxy-5-methyl-
				• • • • • • • • • • • • • • • • • • • •	-			1.

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance Data are for structurally related substance, C.I. Acid Red 14, 1-

Naphthalenesulfonic acid, 4-hydroxy-3-[(4-sulfo-1-

naphthalenyl)azo]-, disodium salt (CAS No. 3567-69-9)

Method Not given

GLP Ambiguous

Year 1993

Contact time (units) 24 hour

Innoculum Activated sludge

Remarks for Test Conditions Screened raw wastewater was used as the influent in three

pilot scale activated sludge biological treatment systems. Each water soluble dye was tested at doses of 1 mg/L for low spike systems and 5 mg/L for high spike systems of influent flow. Before the data collection, dye analytical recovery studies were conducted by dosing the purified dye compound into organic free water, influent wastewater, and mixed liquor. These studies were run in duplicate and each recovery study was repeated at least once to ensure that the dye compound could be extracted. Purified dye standards were analytically prepared from the commercial dye product by repeated recrystallization.

The INF, primary effluent (PE), and ASE were filtered and the filtrate was passed through a column packed with resin. The filter paper and resin were soaked in an ammonia acetonitrile solution and then Soxhlet extracted with ammonia-acetonitrile. The extract was concentrated and brought up to 50 mL volume with a methanol/dimethylformamide solution. The mixed liquor samples were separated into two components, the filtrate or soluble fraction (SOL) and the residue (RES) fraction. The SOL fraction was processed similar to these samples but he resin adsorption step was omitted. All extracted samples were analyzed by HPLC with and ultraviolet-visible detector. Total suspended solids analyses were also performed on the INF. PE, ML, and ASE samples.

All systems were operated for at least three times the solids retention time to ensure acclimation prior to initiation of data collection. All samples were 24 hr. composites made up of 6 grab samples collected every 4 hr. and stored at 4 deg Celsius. Percent recovery as measured: Organic Free Water: 101% at 1 mg/L and 90% at 5 mg/L; Wastewater: 98% at 1mg/L and 97% at 5 mg/L; Mixed Liquor: 88% at 1mg/L and 92% at 5 mg/L Mass Balance Data Summary: Low spike: 116% recovered, 1% adsorbed; High spike: 148% recovered, less than 1%

adsorbed.

Remarks fields for results Since the majority of the test substance was recovered, the

authors assumed that this compound was not biodegraded. The authors based this assumption on preliminary data indicating little or no problems in recovering the compounds from the sample matrix. Additionally the results also indicate that the material was not adsorbed. The authors attributed the high sulfonic acid substitution on the test substance as the reason why the material was not removed by the microbial cells

or cell byproducts and subject to aerobic biodegradation.

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Comparable to guideline study.

Shaul G.M., Holdsworth T.J., Dempsey C.R., and Dostal K.A. Reference

(1990) Fate of water soluble azo dyes in the activated sludge

process. Chemosphere 22, p107-119.

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-Substance Name

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Results

Method Calculated

Classification Not readily biodegradable

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

Reference BIOWIN EPI Suite (2000) US Environmental Protection

Agency.

2.3 FUGACITY

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Air-Water Partition Coefficient

Absorption coefficient 1.13E-18

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Fish-Water Partition Coefficient

Absorption coefficient 0.0491

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Aerosol-Air Partition Coefficient

Absorption coefficient 2.87E+15

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Air

Estimated Distribution and Media Concentration

3.68E-24%

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald

(1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Soil-Water Partition Coefficient

Absorption coefficient 0.0201

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Sediment-Water Partition Coefficient

Absorption coefficient 0.0403

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-
	methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Suspended Sediment-Water Partition Coefficient

Absorption coefficient 0.201

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-
	methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Water

Estimated Distribution and

Media Concentration

Data Qualities Reliabilities

100%

Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt			
CAS No.	25956-17-6			
Model Conditions	25 °C, 100,000 pounds			
Test Type	Environmental Equilibrium Partitioning Model			
Method	Mackay			
Model Used	EQC V 2.70 Level III			
Input Parameters	MW, log Kow, water solubility, MP & VP			
Media	Soil			
Estimated Distribution and	1.64E-14%			
Media Concentration Data Qualities Reliabilities	Reliability code 4. Not assignable.			
Remarks for Data Reliability	Code 4. Calculated.			
References	Trent University (2002) Level III Fugacity-based Environmental Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.			
Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt			
CAS No.	25956-17-6			
Model Conditions	05.00 400.000			
	25 °C, 100,000 pounds			
Test Type	Environmental Equilibrium Partitioning Model			
Test Type Method	•			
•	Environmental Equilibrium Partitioning Model			
Method	Environmental Equilibrium Partitioning Model Mackay			
Method Model Used	Environmental Equilibrium Partitioning Model Mackay EQC V 2.70 Level III			
Method Model Used Input Parameters	Environmental Equilibrium Partitioning Model Mackay EQC V 2.70 Level III MW, log Kow, water solubility, MP & VP			
Method Model Used Input Parameters Media	Environmental Equilibrium Partitioning Model Mackay EQC V 2.70 Level III MW, log Kow, water solubility, MP & VP Sediment			
Method Model Used Input Parameters Media Absorption coefficient	Environmental Equilibrium Partitioning Model Mackay EQC V 2.70 Level III MW, log Kow, water solubility, MP & VP Sediment 0.0403%			

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-
	methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

1.01E-4%

Media Suspended Sediment

Estimated Distribution and Media Concentration

Data Qualities Reliabilities

Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-

methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Model Conditions 25 °C, 100,000 pounds

Test Type Environmental Equilibrium Partitioning Model

Method Mackay

Model Used EQC V 2.70 Level III

Input Parameters MW, log Kow, water solubility, MP & VP

Media Fish

Estimated Distribution and Media Concentration Data Qualities Reliabilities

4.91E-6%

Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

References Trent University (2002) Level III Fugacity-based Environmental

Equilibrium Partitioning Model Version 2.70. Mackay, Donald (1991) Multimedia environmental models: The fugacity approach. Lewis Publications, CRC Press, Boca Raton, FL.

3 ECOTOXICITY

3.1 Acute Toxicity to Fish

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt
CAS No.	25956-17-6
Remarks for Substance	Data are for structurally related substance, 2,2'-(1,2-ethene-diyl)bis(5-amino)-benzenesulfonic acid (CAS No. 81-11-8)
Test Type	Experimental
GLP	Ambiguous
Year	Not given
Species/Strain/Supplier	Fish
Exposure Period	48 hour
Endpoint value	LC50 = 200 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Remarks for Data Reliability	Code 4.Only secondary literature (review, tables, books, etc.).
Reference	Greim H., Ahlers J., Bias R., Broecker B., Hollander H., Gelbke H.P., Klimisch H., Mangelsdorf I., Paetz A., Schone N., Stropp G., Vogel R., Weber C., Ziegler-Skylakakis K., and Bayer E. (1994) Toxicity and ecotoxicity of sulfonic acids: structure-activity relationship. Chemosphere, 28, 2203-2236.
Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt
CAS No.	25956-17-6
Remarks for Substance	Data are for structurally related substance, 2,2'-(1,2-ethene-

diyl)bis(5-amino)-benzenesulfonic acid, disodium salt (CAS No. 7336-20-1)

Test Type Experimental

GLP Ambiguous

Year Not given

Species/Strain/Supplier Fish

Exposure Period 72 hour

Endpoint value LC50 greater than 1000 mg/L

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4.Only secondary literature (review, tables, books, etc.).

Reference Greim H., Ahlers J., Bias R., Broecker B., Hollander H., Gelbke

H.P., Klimisch H., Mangelsdorf I., Paetz A., Schone N., Stropp G., Vogel R., Weber C., Ziegler-Skylakakis K., and Bayer E. (1994) Toxicity and ecotoxicity of sulfonic acids: structure-

activity relationship. Chemosphere, 28, 2203-2236.

Substance Name

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance Data are for structurally related substance, 2,2'-(1,2-ethene-

diyl)bis(5-amino)-benzenesulfonic acid, dipotassium salt (CAS

No. 78447-91-3)

Test Type Experimental

GLP Ambiguous

Year Not given

Species/Strain/Supplier Fish

Exposure Period 96 hour

Endpoint value LC50 greater than 10000 mg/L

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4.Only secondary literature (review, tables, books, etc.).

Reference Greim H., Ahlers J., Bias R., Broecker B., Hollander H., Gelbke

H.P., Klimisch H., Mangelsdorf I., Paetz A., Schone N., Stropp G., Vogel R., Weber C., Ziegler-Skylakakis K., and Bayer E. (1994) Toxicity and ecotoxicity of sulfonic acids: structure-

activity relationship. Chemosphere, 28, 2203-2236.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Method/guideline ECOSAR

Test Type Calculated

Species/Strain/Supplier Fish

Exposure Period 96 hour

Remarks for Test Conditions Input parameters: Melting point, 350 °C, Water solubility,

220,000 mg/L at 25 deg C

Endpoint value LC50 = 2714 mg/L

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

Reference ECOSAR EPI Suite (2000) U.S. Environmental Protection

Agency (Nabholz V. and G. Cash, 1998).

3.2 Acute Toxicity to Aquatic Invertebrates

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt
CAS No.	25956-17-6
Remarks for Substance	Data are for structurally related substance 2,2'-(1,2-ethenediyl)bis(5-amino)-benzenesulfonic acid, disodium salt (CAS No. 7336-20-1)
Test Type	Experimental
Species/Strain/Supplier	Daphnia magna
Test Details	24 hour
EC50, EL50, LC0, at 24,48 hours	EC50 = 100 mg/L
Data Qualities Reliabilities	Reliability code 4. Not assignable.
Data Reliability Remarks	Code 4.Only secondary literature (review, tables, books, etc.).
Reference	Griem H., Ahlers J., Bias R., Broecker B., Hollander H., Gelbke H.P., Klimisch H., Mangelsdorf I., Paetz A., Schone N., Stropp G., Vogel R., Weber C., Ziegler-Skylakakis K., and Bayer E. (1994) Toxicity and ecotoxicity of sulfonic acids: structure-activity relationship. Chemosphere, 28, 2203-2236.
Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt
CAS No.	25956-17-6

Method/guideline ECOSAR

Test Type Calculated

Species/Strain/Supplier Daphnia magna

Test Details 48 hours

Remarks for Test Conditions Input parameters: Melting point, 350 °C, Water solubility,

220,000 mg/L at 25 deg C

EC50, EL50, LC0, at 24,48

hours

LC50 = 295 mg/L

Data Qualities Reliabilities Reliability code 4. Not assignable.

Data Reliability Remarks Code 4. Calculated.

Reference ECOSAR EPI Suite (2000) U.S. Environmental Protection

Agency (Nabholz V. and G. Cash, 1998).

3.3 ACUTE TOXICITY TO AQUATIC PLANTS

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4 cultophonyl)azal disadium salt

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

substituted azo dye.

Test Type Experimental

GLP Ambiguous

Year 1996

Species/Strain/Supplier Green algae, Selenastrum capricornutum

Exposure Period 96 hour

Remarks for Test Conditions Algal chronic toxicity test were performed according the method

of EPA, 1988. Three replicates were performed for each dye at a nominal concentration of 1 mg/l for the active colorant. One ml of dye stock solution was added to 50 mg/l of algal assay medium in 125 ml Erlenmeyer flasks. S. capricornutum in continuous culture provided the initial innoculum (10,000 algal cells/ml). The cells were incubated in the solution for 96 hours. The diluent and negative control were algal assay medium. AAM was prepared by adding 1 ml from each of five stock

solutions to 900 ml of deionized water. After spiking, the total volume was brought to 1 liter with deionized water. Population growth was used to establish potential toxicity. If the dye inhibited algal growth by more than 50% of that of the negative controls, a definitive test using several dilutions of the dye was performed to allow for determination of an EC50 concentration.

Endpoint value Average yield: 36.6% with 95% C.I. (34.9-38.4).

Biological observations 26.4% stimulation of population growth compared to control.

Control response satisfactory?

Yes

Appropriate statistical

evaluations?

Yes, Dunnett's test

Remarks fields for results
Not statistically significant.

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 1. Comparable to guideline study.

Reference Greene J. C. and Baughman G.L. (1996) Effects of 46 dyes on

population-growth of fresh-water green-alga *selenastrum-capricornutum*. Textile Chemist And Colorist, 28, 23-30. Green J.D. et al. (1988) Protocols for short term toxicity screening of hazardous waste sites. Report to EPA 600/3-88-029. U.S. Environmental Protection Agency. Corvallis, Oregon.

Substance Name

2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Method/guideline ECOSAR

Test Type Calculated

Species/Strain/Supplier Green algae

Exposure Period 96 hour

Remarks for Test Conditions Input parameters: Melting point, 350 °C, Water solubility -

220,000 mg/L at 25 deg C

Endpoint value EC50 = 44,524 mg/L

Data Qualities Reliabilities Reliability code 4. Not assignable.

Remarks for Data Reliability Code 4. Calculated.

Reference ECOSAR EPI Suite (2000) US Environmental Protection

Agency (Nabholz V. and G. Cash, 1998).

4 HUMAN HEALTH TOXICITY

4.1 ACUTE TOXICITY

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4-sulfonhenyl)azol- disodium salt

CAS No. 25956-17-6

Remarks for Substance FD&C Red No. 40; purity not given; dark red in color

Method/guideline Not given

Test Type Acute Oral LD50

GLP No

Year 1965

Species/strain Rat/Sprague-Dawley albino

Sex Male and Female

of animals per sex per

dose

5 male and 5 female

Vehicle Water

Route of Administration Oral-Gavage

Remarks for Test Condition Six groups of five male and five female Sprague-Dawley rats

each were administered the test substance in a 10% weight/volume solution. The dosage levels tested were 215, 464, 1000, 2150, 4640, and 10,000 mg/kg bw. The animals were fasted for 3-4 hours prior to dosing. Following dosing, the animals were housed in metal cages suspended above the droppings. Food and water were available *ad libitum*.

Observations were made immediately following dosing, at 1, 4, 24, 48 hours and once daily thereafter up to 14 days. Following the observation period, the animals were weighed, sacrificed by

cerebral concussion and necropsied.

Greater than 10,000 mg/kg bw

Value LD50 or LC50 with

confidence limits

Number of deaths at each

dose level

There were no deaths at any dose level tested.

Remarks for results

Clinical observations were normal with the exception of redcolored feces in both sexes at all dose levels and red-colored urine at the three highest dose levels in the female animals.

Conclusion remarks

The acute LD50 was determined to be greater than 10,000

mg/kg bw/d for adult male and female Sprague-Dawley albino

rats.

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability

Code 2. Basic data given: comparable to guidelines/standards.

References

Hazelton Laboratories, Inc. (1965a) Acute oral administrationrats. Five experimental non-toxic red colors. Unpublished

Report No. 165-114.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance FD&C Red No. 40; purity not given; dark red in color

Method/guideline Not given

Test Type Acute Oral LD50

GLP No

Year 1965

Species/strain Dog/Mongrel

Sex Male

of animals per sex per

dose

2 males

Vehicle Water

Route of Administration Oral-Gavage

Remarks for Test Conditions One groups of two male Mongrel dogs was administered the

test substance in an aqueous solution at a dose level of 5 g/kg bw. Two concurrent control animals receiving 300 ml of water each were also maintained. Each test animal was individually

housed. Food and water were available ad libitum.

Observations were made immediately following dosing and daily thereafter for 7 days. Following the observation period, the animals were weighed, sacrificed and necropsied. Necropsies

were not performed on control animals. Greater than 5,000 mg/kg bw

Value LD50 or LC50 with confidence limits

Number of deaths at each

number of deaths at each

dose level

Remarks for results

There were no deaths at the dose level tested (5000 mg/kg

bw).
Red diarrhea was observed 30 minutes following dosing in one

animal, which was followed by emesis. Red urine was reported for the other animal. Red stools were reported for both dogs one day following dosing. From the third day until the seventh day, both animals appeared normal with respect to appearance, behavior, appetite and elimination. Gross necropsy revealed fibrotic changes and decreased weight in a kidney of one test animal. This finding was not considered treatment-related but was rather considered to be a chronic lesion. The spleen also appeared enlarged in this test animal. In the other test animal, hookworms were observed in the

gastrointestinal tract.

Conclusion remarks The acute LD50 was determined to be greater than 5,000

mg/kg bw/d for male Mongrel dogs.

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Hazelton Laboratories, Inc. (1965b) Acute oral administration-

dogs. Five experimental non-toxic red colors. Unpublished

Report.

4.2 GENETIC TOXICITY

4.2.1 In vitro Genotoxicity

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4 authoritant and landium ant

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance Purity not given; red powder.

Method/guideline Ames, McCann and Yamasaki (1975) Plate Test (Overlay

method)

Test Type Reverse mutation

System of Testing Bacterial

GLP Ambiguous

Year 1983

Species/Strain Salmonella typhimurium TA98, TA1535, TA1537;

Saccharomyces cerevisiae strain D4

Metabolic Activation Rat liver microsome fraction S9 from Aroclor induced rats

Doses/Concentration 0.625, 1.25, 2.5, 5.0% or 10, 100, 1000, or 5000 micrograms

per plate

Statistical Methods Not given

Remarks for Test Conditions Toxicity tests were conducted to identify the 12.5%, 25% and

50% killing doses. If no toxicity was found, a maximum dose of 5% was used as the highest dose concentration. The same doses were used for both activation and non-activation assays. Approximately 10^9 cells from a log phase culture of each indicator strain were added to test tubes containing 2.0 ml of molter agar supplemented with biotin and a trace of histidine.

For tests with activation, the rat liver 9000 x g tissue

supernatant and required cofactors were added to the overlay tubes. The four dose levels of the test substance were added to the overlay tubes, followed by mixing and pouring over minimal agar plates. The plates were then incubated for 48-72 hours at 37 deg Celsius and scored for colonies. Positive and negative (solvent only) controls were run with each assay. Positive

controls for the non-activation assays were

ethylmethanesulfonate (EMS); methylnitrosoguanidine (MNNG); 2-nitrofluorene (NF); quinacrine mustard (QM). Postive controls with activation included 2-anthramine (ANTH);

2-acetylaminofluorene (AAF); 8-aminoquinoline (AMQ);

dimethylnitrosamine (DMNA).

Results The maximum dose of 5% was used, since 50% survival was

not determined. Slight toxic effects noted at 5%.

Cytotoxic concentration Slight toxic effects at 5%.

Genotoxic Effects Negative at all concentration levels.

Appropriate statistical

evaluations?

None given

Remarks for results The substance was determined to be soluble.

Conclusion Remarks The test substance did not exhibit genotoxic activity with or

without metabolic activation in the AMES assay using SAL

TA98, TA1535 or TA1537 plate overlay method.

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Acceptable, well-documented publication/study report,

which meets basic scientific principles.

References Brusick D. (1976) Mutagenicity evaluation of NTR-Z-4576.

Unpublished report.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

4-sulfophenyl)azol-, disodium salt

CAS No. 25956-17-6

Remarks for Substance Purity not given

Method/guideline Ames test

Test Type Reverse mutation

System of Testing Bacterial

GLP No

Year 1979

Species/Strain Salmonella typhimurium TA1535, TA 1537, TA98, TA100

Metabolic Activation Rat liver microsome fraction S9 from Aroclor induced rats

Doses/Concentration 10-250 mg/plate

Statistical Methods Not given

considered positive if 2 fold increase in revertants was observed. Positive controls included 9-aminoacridine; 2-

aminoflourine; N-methyl-N-nitrosoguanidine.

Results Negative

Cytotoxic concentration Not given

Genotoxic Effects Negative

Appropriate statistical

evaluations?

None given

Remarks for Results Negative

Conclusion Remarks No evidence of genotoxicity was reported.

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Basic data given: comparable to guidelines/standards.

References Muzzall J.M. and Cook W.I. (1979) Mutagenicity test of dyes

used in cosmetics with the Salmonella/mammalian microsome

test. Mutations Research 67, 1-8.a

4.2.2 In vivo Genotoxicity

	4-sulfophenyl)azo]-, disodium salt
Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

CAS No. 25956-17-6

Yellow 6, 6-hydroxy-5-[(4-sulfophenyl)azo]-2-

napthalenesulfonic

acid, disodium salt (CAS No. 2783-94-0).

Method/guideline Rodent Micronucleus Test

GLP Ambiguous

Year 1991

Species/Strain Rat/PVG

Sex Male

Route of Administration Oral-Gavage

Doses/Concentration 10 ml/kg bw

Exposure Period Single dose

Remarks for Test Conditions Male PVG rats received a single oral dose of 500, or 1000

mg/kg of sunset yellow 6. Bone marrow samples were taken at

24 and 48 hours later.

Genotoxic effects No significant increase in the frequency of micronucleated

polychromatic erythrocytes at either time point in either species

and there was no effect on the % PE (polychromatic

erthyrocytes).

Appropriate statistical

evaluations?

Remarks for results No

No effects.

Yes

Data Qualities Reliabilities Reliability code 2. Reliable with restriction.

Remarks for Data Reliability Code 2. Acceptable, well-documented publication/study report,

which meets basic scientific principles.

References Westmoreland C. and Gatehouse D.G. (1991) The differential

clastogenicity of Solvent Yellow 14 and FD & C Yellow No. 6 in vivo in the rodent micronucleus test (observations on species

and tissue specificity). Carcinogenesis 12 (8), 1403-8.

4.3 REPEATED DOSE TOXICITY

Substance Name2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance FD&C Red No. 40; 88% purity

Method/guideline Lifetime Toxicity/Carcinogenicity Study

GLP Ambiguous

Year 1991

Species/strain Rat/Sprague-Dawley

Sex Male and Female

Route of Administration Oral-Diet

Doses/concentration Levels 0.37, 1.39 or 5.19%

Exposure Period 118 (males) or 121 weeks (females)

Frequency of Treatment

Daily

Control Group

Yes

Remarks for Test Conditions

In a Lifetime Toxicity/Carcinogenicity Study, FD & C Red 40 was provided in the diet as an admixture to Sprague-Dawley rats. In the in utero phase, 240 male and female rats were randomly assigned (30/group) to the control, low dose (0.37%). mid-dose (1.39%) or high dose (5.19%) groups, providing daily intake levels of 180, 701 or 2829 mg/kg bw/d for males and 228, 901 or 3604 mg/kg bw/d for females. These parental (P1) rats received the test material one week prior to mating, during the three-week mating period and during the gestation and lactation periods. The offspring of these animals were randomly selected and put into groups of fifty male and female weanling rats each. These groups were administered the test substance in the diet of the male animals for 118 weeks and the diet of female animals for 121 weeks at levels of 0, 0.37, 1.39 to 5.19 % corresponding to the dietary levels used in the in utero phase. Parameters included survival, clinical signs, body weight and food consumption, gross and microscopic pathology. Gross necropsies were performed on all animals dying during the study, all animals found in a moribund condition, and all animals killed at study termination. Complete histological examinations were performed on all animals in both the control and high-dose groups. The tissues examined histologically included: brain, pituitary, thoracic spinal cord, eyes, esophagus, thyroid, thymus, heart, lungs, liver, spleen, pancreas, stomach, small and large intestine, mesenteric lymph node, kidneys, adrenal, urinary bladder, uterus, prostate, ovaries, testes with epididymides, seminal vesicles, skin, rib junction, bone marrow, nerve with muscle, and any tissue masses or lesions. Histological examination was also performed on animals from any group with observable masses or lesions. If a potential effect was seen recurrently in a tissue, than that tissue was examined in all animals.

NOAEL(NOEL)

LOAEL(LOEL)

Actual dose received by dose level and sex Toxic Response/effects by Dose Level 5.19% or 2829 mg/kg bw/d (males); 1.39% or 901 mg/kg bw/d (females)

Greater than 5.19% or 2829 mg/kg bw/d (males); 5.19% or 3604 mg/kg bw/d (females)

180, 701 or 2829 mg/kg bw/d (males); 228, 901 or 3604 mg/kg bw/d (females)

Food consumption was elevated among high dose males and females, but was not statistically significant. Red-tinted fur was reported among all treated animals, and red-tinted feces was reported among mid- and high-dose male and females. Group mean body weights of treated males and females were decreased compared to control animals at study termination, with the exception of mid-dose treated male rats, which experienced an increase in mean body weight. However, the decrease in mean body weight was only statistically significant in female rats at the high dose level (3604 mg/kg bw/d). Clinical chemistry and urinalysis parameters revealed no treatment related effects. Histopathological examination revealed lesions in both control and treated animals at similar prevalence, and thus not attributed to test substance administration.

Appropriate statistical

evaluations?

Yes

Conclusion Remarks

No biologically significant adverse effects were reported following administration of FD&C Red 40, with the exception of

decrease mean body weights for high-dose female rats at study termination. The authors attributed this effect to the large amount of non-nutritive material in the diet at the intake level.

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Comparable to guideline study.

References Borzelleca J.F., Olson J.W. and Reno F.E. (1991a) Lifetime

toxicity/ carcinogenicity studies of FD&C Red No. 40 (Allura Red) in Sprague Dawley Rats. Food and Chemical Toxicology,

27, 701-705.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Test Substance FD&C Red No. 40; 88% purity

Method/guideline Lifetime Toxicity/Carcinogenicity Study

GLP Ambiguous

Year 1991

Species/strain Mice\Charles River CD1 (study A) and outbred CD-1 (study B)

Sex Male and Female

Route of Administration Oral-Diet

Doses/concentration Levels 0.37, 1.39 or 5.19%

Exposure Period 104 weeks (Study A) or 109 weeks (Study B)

Frequency of Treatment Daily

Control Group Yes

Remarks for Test Conditions In the in utero phase, 50 male and female mice each (study A)

or 70 male and female mice each (study B) were randomly assigned to the control, low dose (0.37%), mid-dose (1.39%) or high dose (5.19%) groups, providing daily intake levels of 507, 1877 or 7422 mg/kg bw/d for males and 577, 2043 or 8304 mg/kg bw/d for females (study A) and 492, 1821, or 7318 mg/kg bw/d (males) and 526, 2057 or 8356 mg/kg bw/d (females) (study B). These Fo mice received the test material one week prior to mating, during the three week mating period and during gestation and lactation periods. Groups of fifty male

and female weanling Charles River mice each were

administered the test substance in the diet of study A animals for 104 weeks and the diet of study B animals for 109 weeks at levels of 0, 0.37, 1.39 or 5.19 %. These animals were the Fo

offspring of parental mice (P1), which were treated at the corresponding levels. Study A had one control group while study B had two control groups. Parameters included survival, clinical signs, body weight and food consumption, gross and microscopic pathology. Gross necropsies were performed on all animals dying during the study, all animals found in a moribund condition, and all animals killed at study termination. Complete histology was conducted on all mice from all groups in study A and on 10/sex/group for the two control groups and the highest-dose group from study B. The tissues examined histologically included: brain, pituitary, thoracic spinal cord, eyes, esophagus, thyroid, thymus, heart, lungs, liver, spleen, pancreas, stomach, small and large intestine, mammary glands (study B only), mesenteric lymph node, kidneys, adrenal. urinary bladder, uterus, prostate, ovaries, testes with epididymides, seminal vesicles, skin, rib junction, bone marrow. nerve with muscle, and any tissue masses or lesions. Greater than 5.19%

NOAEL(NOEL)

LOAEL(LOEL)

Actual dose received by dose level and sex

Toxic Response/effects by Dose Level
Appropriate statistical evaluations?
Remarks for Results

Not determined

507, 1877 or 7422 mg/kg bw/d for males and 577, 2043 or 8304 mg/kg bw/d for females (study A) and 492, 1821, or 7318 mg/kg bw/d (males) and 526, 2057 or 8356 mg/kg bw/d (females) (study B).

No treatment -related effects were observed for any parameter evaluated at any dose level.

Yes.

No treatment-related effects were reported on survival. The authors reported decreased food consumption among the midand high-dose females for wk 62-106 in study B. However, no consistent statistically significant effects on food consumption were reported in either study. Localized alopecia, labored respiration, colored hair coat, lacrimation and thinness were reported in similar incidences in both control and treated mice at all dose levels. Distended abdomens were noted in both midand high-dose females, while palpable masses were reported in control and treated groups at a similar incidence. Hematological and clinical chemistry parameters revealed few differences among treated and control groups. No significant gross pathological changes were reported among treated groups compared to control groups. An increase in absolute and relative thyroid weights in study B in the high-dose males and females was reported but the significance was questioned because there was no accompanying histopathology, and were not dose-dependent and were species-specific.

The authors also reported an earlier appearance of lymphatic lymphomas among treated groups in study A compared to control groups. No increases in incidence or appearance of lymphocytic lymphomas was reported in study B. However, statistical analyses of the data revealed no statistical significance in the finding of an apparent acceleration of lymphocytic lymphomas development.

Conclusion Remarks No treatment-related adverse effects were reported at any dose

level following lifetime administration of FD & C Red 40 to male

and female mice.

The second study, study B, conducted using a different strain of mouse to further investigate if FD&C Red 40 had an effect on the appearance of lymphocytic lymphomas, revealed no relationship between the incidence of lymphocytic lymphomas

and FD&C Red 40.

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Comparable to guideline study.

References Borzelleca J.F., Olson J.W. and Reno F.E. (1991b) Lifetime

toxicity/ carcinogenicity studies of FD&C Red No. 40 (Allura Red) in mice. Food and Chemical Toxicology, 29, 313-319.

4.4 DEVELOPMENTAL TOXICITY

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4 aultanhanyl\azal diaadium aalt

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Method/guideline FDA Teratology Study

GLP Yes

Year 1989

Species/strain Rat/Osborne-Mendel (FDA strain)

Sex Female

Route of Administration Oral-drinking water

Duration of Test 20 days

Doses/concentration Levels 0, 0.2, 0.4 or 0.7%

Exposure Period 20 days

Frequency of Treatment ad libitum

Control Group and

Treatment

Yes

Remarks for Test Conditions

Four groups of female Osborne-Mendel (FDA strain) rats (40-41 per group) were administered FD & C Red 40 in the drinking water at intake levels of 0, 0.2, 0.4 or 0.7% for the first 20 days of gestation. On day 20, the animals were examined for gross abnormalities followed by euthanization. Caesarean sections were performed. The uterus was examined for presence and position of resorption sites and fetuses, number of corpora lutea and implantation sites. All live fetuses were promptly weighed, sexed, and examined. Crown-rump lengths were measured. Fetuses were divided and assigned to skeletal or soft tissue examination.

NOAEL(NOEL) maternal

toxicity

LOAEL(LOEL) maternal

toxicity

NOAEL (NOEL)

developmental toxicity

LOAEL (LOEL)

developmental toxicity Actual dose received by dose level and sex

Maternal data with dose level

Fetal Data with Dose Level

Appropriate statistical evaluations? Remarks for results

Data Qualities Reliabilities

Remarks for Data Reliability

Not determined

.7% or 939.29 mg/kg bw/d

273.58 mg/kg bw/d

545.68 mg/kg bw/d

0, 273.58, 545.68 or 939.29 mg/kg bw/d

No clinical findings were reported and no deaths occurred during treatment. Mean fluid consumption was significantly increased in animals at the 0.2 and 0.4% intake levels but only on days 14-20. Because fluid consumption was not increased at the 0.7% level, the findings were not considered biologically

significant. No other effects were reported.

A significant increase in the incidence of litters containing fetuses with missing sternebrae occurred in the 0.4% group, but not in the group receiving 0.7%. No dose related increases were reported for any sternebral variations. The number of fetuses with at least one type of sternebral variations was greater in all treated groups, but only significantly greater in the 0.4 and 0.7% groups. The percentage of total fetuses with at least one sternebral variation was greater in all of the treated groups compared to the control group, but the differences were not significant. The number of fetuses with more than one skeletal variation were similar among treated and control groups. The incidence of reduced ossification of the hyoid bone was significantly increased at the 0.7% intake level. Significant dose related increases were reported at the highest intake level for the average number of fetuses per litter with at least two skeletal variations and the number of litters containing them. Yes, ANOVA, Fisher's Exact Test, t-test.

The authors questioned the biological significance of the reduced ossification of the hyoid bone, given the lack of effect seen in a gavage study using higher dose levels. The increased incidence was also just outside that found in the historical controls, and the control group was noted as having a lower incidence compared to the historical controls.

Reliability code 1. Reliable without restriction.

Code 1. Guideline study.

References Collins T., Black T.N., Welsch J.J., and Brown L.H. (1989a)

Study of the teratogenic potential of FD & C Red No. 40 when given in drinking water. Toxicology and Industrial Health 5,

937-948.

Substance Name 2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-

4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Method/guideline FDA Teratology Study

GLP Yes

Year 1989

Species/strain Rat/Osborne-Mendel (FDA strain)

Sex Female

Route of Administration Oral-Gavage

Duration of Test 19 days

Doses/concentration Levels 0, 30, 75, 150, 300, 600 or 1000 mg/kg bw/d

Exposure Period 19 days

Frequency of Treatment Daily

Control Group and

Treatment

Remarks for Test Conditions

Yes

Four groups of female Osborne-Mendel (FDA strain) rats (42-43 per group) were administered FD & C Red 40 via gavage at dose levels of 0, 30, 75, 150, 300, 600 or 1000 mg/kg bw/d for the first 19 days of gestation. On day 19, the animals were examined for gross abnormalities followed by euthanization. Caesarean sections were performed. The uterus was examined for presence and position of resorption sites and fetuses, number of corpora lutea and implantation sites. All live fetuses were promptly weighed, sexed, and examined. Crown-rump lengths were measured. Fetuses were divided and assigned to

skeletal or soft tissue examination.

NOAEL(NOEL) maternal

toxicity

LOAEL(LOEL) maternal

toxicity

NOAEL (NOEL)

developmental toxicity

LOAEL (LOEL)

developmental taxiait

developmental toxicity Appropriate statistical

evaluations?

Actual dose received by dose level and sex

Maternal data with dose

level

1000 mg/kg bw/d

Not determined

1000 mg/kg bw/d

Not determined

Yes, ANOVA, Fisher's Exact Test, t-test.

0, 30, 75, 150, 300, 600 or 1000 mg/kg bw/d

No clinical findings were reported and no deaths occurred

during treatment. No other dose related findings were reported.

Fetal Data with Dose Level The only significant skeletal anomaly found was an increase in

14th rib buds at the 300 mg/kg bw/d dose level but was not seen at the higher dose levels. No other soft-tissue or

sternebral variations were reported.

Conclusion remarks The NOAEL's for maternal and fetal toxicity were 1000 mg/kg

bw/d.

Data Qualities Reliabilities Reliability code 1. Reliable without restriction.

Remarks for Data Reliability Code 1. Guideline study.

References Collins T., Black T.N., Welsch J.J., and Brown L.H. (1989b)

Study of the teratogenic potential of FD & C Red No. 40 when given by gavage to rats. Fd. Chem. Toxic. Vol 27, pp 707-713.

4.5 REPRODUCTIVE TOXICITY

Substance Name	2-Naphthalenesulfonic acid, 6-hydroxy-5-[(2-methoxy-5-methyl-
	4-sulfophenyl)azo]-, disodium salt

CAS No. 25956-17-6

Remarks for Substance FD&C Red No. 40; fine dark red powders without noticeable

odor

Method/guideline Not given

Test Type Two generation reproductive study

GLP Ambiguous

Year 1969

Species/strain Rat/Charles River Caesarean albino

Sex Male and Female

Route of Administration Oral-Diet

Duration of TestTwo parental generations and two two-litter filial generations

Doses/concentration Levels 3700, 13,900 and 51,900 ppm

Premating Exposure period

for males

27 weeks

Premating Exposure period

for females

27 weeks

Frequency of Treatment

Daily

Control Group and Treatment Remarks for Test Conditions

Yes, basal diet

Groups of male (10) and female (20) Charles River rats were administered FD&C Red No. 40 in the diet at 0, 3700, 13,900, or 51,900 ppm for 27 weeks prior to initiation of the first breeding phase. These P1 parental generations were individually housed. Clinical observations included food consumption, appearance, individual body weights and behavior and were made weekly. The F1A weanling rats designated P2 generation were kept 4-5 to a cage according to sex and maintained on the same concentration level as their parents until reaching maturity.

During the breeding phase of the P1 generation, two females and one male were placed in a breeding cage. At weekly intervals during the mating period, the males were rotated among the females in each group. Following mating, the females were placed in individual cages to produce the first (FIA) litters. Twenty-four hours following the birth of the pups the first litters (FIA) were arbitrarily reduced to 8 maximum per mother. The number of conceptions, number of litters, live births, stillbirths, size of natural and nursing litters, deaths during the period of lactation, and number of pups weaned were recorded. The body weights of each pup were recorded at 24 hours and at weaning. Gross signs of toxicity were monitored. After 21-days of nursing, random pups were sacrificed and gross necropsies performed. Twenty-four females and twelve males remaining from each test group and control group were selected at random and designated the P2 generation. Following the weaning of the F1A animals, the P1 generation was remated to produce their second litters referred to as F1B, according to the procedures described above.

The P2 generation was housed 4-5 per cage and was maintained on the same dietary levels as their parents. The procedures outlined above for the P1 generation were maintained for the P2 generation. The litters of the P2 animals were referred to as the F2A litters. Body weights of the F2A pups were monitored 24 hours following the birth and at weaning. Gross signs of toxicity were recorded. Following a 21-day nursing period, all pups were weaned and sacrificed. One week following the weaning period of the F2A litter, the P2 generation was remated to produce their second litters (F2B). Two females were placed in a cage with a male from the corresponding dose group. Males were rotated weekly, and females were examined daily for presence of spermatozoa for a maximum of 21 consecutive days. The first day that sperm were observed was designated as day 0 of gestation. The females were then placed in individual cages. Half of the females (12) were sacrificed on day 19 or 20 of gestation and Caesarean sections were performed. Observations included number and placement of implantation sites, resorption sites, and live and dead fetuses; individual fetal weight and length (crown to rump), and external fetal anatomical structure. Gross necropsies were performed on each female including examination of uterus and visceral structures. The remaining 12 females were allowed to litter normally. The fetuses of both females delivering normally and via Caesarean section were

necropsied.

NOAEL(NOEL) 13,900 ppm

LOAEL(LOEL) 51,900 ppm

Actual dose received by dose level and sex Parental data and F1 as

Not given

appropriate

Fertility indices for the control and test animals of both F1A and F1B were considered low. The authors attributed this to the advanced age of the animals upon mating. The fertility index of the 3700 ppm test group in the F2A breeding cycle as well as the 3700 and 51900 ppm test groups in the F2B breeding cycle were reported to be low in comparison to control animals and historical control data.

Offspring toxicity F1 and F2

Growth suppression characterized as slight was also reported for the low-level F1B pups, and the high-level F1A and F1B pups and the F2A and F2B breeding cycles when compared with controls. All other measured parameters were comparable to controls in each generation and among the two filial generations. The authors concluded that FD&C Red 40 caused meaningful growth suppression in the pups whose parents received the high level diets.

Appropriate statistical evaluations?

Not given

Conclusion remarks

The authors reported a NOAEL for reproductive toxicity following administration of FD&C Red 40 as 13,900 ppm.

Data Qualities Reliabilities

Reliability code 1. Reliable without restriction.

Remarks for Data Reliability

Code 1. Comparable to guideline study.

References

Hazelton Laboratories Inc. (1969) Two-generation reproductive study in rats. Red Z4576 (FD&C Red 40). Unpublished report 165-125.